

## Memorial Sloan Kettering Cancer Center

# Transforming Risk Management: Technological Evolution of MSK's **Clinical Research Quality Assurance Program**

BACKGROUND: In 2017, a newly restructured Clinical Research Quality Assurance (CRQA) unit at Memorial Sloan Kettering Cancer Center (MSK) was initiated to substantially improve risk management of the clinical research enterprise. At the time, the unit used systems to collect quality assurance data on clinical research in various formats and platforms that were not originally intended to be interconnected and did not follow symmetrical connection points useful in generating a comprehensive overview of clinical research programs. CRQA's strategy was to restructure and strengthen existing systems, leading to dramatic steps in efficiency and effectiveness while producing an improved employee and customer experience.

### GOALS

CRQA set out to consolidate and improve existing systems to facilitate the extraction of useful data for analyses, while reducing system maintenance. This resulted in restructuring these data collection systems to streamline the collection process and reduce variability while improving report outcomes and accountability. In turn, processes and guidelines required standardization for a comprehensive integration. The overall goal of restructuring was to improve the retrieval and analysis of data to further mitigate institutional risks and identify clinical research process improvement opportunities.



#### **METHODS**



#### **Standardization**

Standardized research-related deficiency definitions and other tools were developed for consistency across CRQA



#### **Enhancements**

Protocol Information Management System (PIMS), MSK's main clinical research information technology system, was enhanced from passive to active



#### **Dashboards**

Developed dashboards for real-time and direct access of data for clinical research departments and CRQA



Interconnections Systems and resources were centralized through data management in PIMS



#### **Data Extraction** Streamlined data extraction were enabled from enhanced systems for better analysis and reporting

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DATA EXTRACTION

RESULTS

The original strategy for PIMS was to transition it from a passive tool to an active tool. In an innovative manner, CRQA went one step further and transitioned the system into an interactive tool accessible not only to CRQA but also to clinical research departments.

The PIMS transformation has since shown to increase compliance, communication, transparency, data access, and collaborative relationships between CRQA and clinical research departments.

The accessibility of real-time data through the development of dashboards, enables CRQA and clinical research departments early identification of deficiencies and trends, while simultaneously identifying potential process improvements.

The dashboards have demonstrated to be powerful tools in highlighting gaps, trends, outstanding regulatory tasks and keeping clinical research departments abreast of these issues.

## **COMCLUSION AND FUTURE DIRECTIONS**

Achievement of these successful improvements required time commitment and collaboration with subject matter experts.

Most of these initiatives needed to be done sequentially to avoid errors and permit testing.

The next goal is to interconnect the data of all CRQA, which consists of four separate programs, into a comprehensive dashboard that can provide a bird's-eye view of all clinical research activities and can forecast and prevent major deficiencies across MSK's clinical research portfolio.

Additionally, we will explore inclusion of automation and advanced analytics into our systems to augment and magnify the impact of process redesign, further enhancing both the effectiveness and efficiency of CRQA's risk management for MSK's clinical research enterprise.